ORIGINAL RESEARCH

Genetic Characteristics of Novel Inc_{pSE5381-aadB} Plasmids, Integrative and Mobilizable Elements, and Integrative and Conjugative Elements in *Pseudomonas aeruginosa*

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Purpose: *Pseudomonas aeruginosa* is a common causative bacteria in nosocomial infections. This study aims to describe the structure and evolutionary characteristics of mobile genetic elements (MGEs) carrying antibiotic resistance genes (ARGs) from *P. aeruginosa* and to conduct bioinformatics and comparative genomic analysis to provide a deeper understanding of the genetic characteristics and diversity of MGEs in *P. aeruginosa*.

Methods: Fifteen clinical isolates of *P. aeruginosa* from China were collected and sequenced in this study, and 15 novel MGEs were identified. Together with four MGEs from GenBank, a total of 19 MGEs were used to perform detailed modular structure dissection and sequence comparison. Then, the biological experiments were carried out to verify the biological characteristics of these isolates and MEGs. **Results:** The novel MGEs identified in this study displayed diversification in modular structures, which showed complex mosaic natures. The seven types of 19 MGEs included in this study were divided into three groups: i) novel MGEs (firstly identified in this study): four $Inc_{pSE5381-aadB}$ plasmids and three Tn7495-related integrative and mobilizable elements (IMEs); ii) newly defined MGEs (firstly designated in this study, but with previously determined sequences): four Tn7665-related IMEs; iii) novel transposons with reference prototypes identified in this study: two Tn6417-related integrative and conjugative elements (ICEs), two IS-based transposition units, two Tn501-related unit transposons, two Tn1403-related unit transposons. At least 36 ARGs involved in resistance to 11 different classes of antimicrobials and heavy metals were identified. Additionally, three novel bla_{OXA} variants were identified. Antimicrobial susceptibility testing showed that these variants were resistant to some β -lactamase antibiotics and $bla_{OXA-1204}$ was additionally resistant to cephalosporins.

Conclusion: The continuous evolution of ARG-carrying MGEs during transmission, leading to the emergence of novel MGEs or ARGs, which facilitates the spread of antibiotic resistance in *P. aeruginosa* and enhances the diversity of transmission modes of bacterial resistance.

Keywords: Pseudomonas aeruginosa, genome sequencing, mobile genetic elements, antimicrobial resistance, OXA

Introduction

Pseudomonas aeruginosa is a common causative Gram-negative bacteria in nosocomial infections, which are often associated with acute and chronic infections in immunocompromised patients (eg post-operative or severe burn patients).¹ The increasing incidence and antibiotic treatment of *P. aeruginosa* infection has made the species resistant to multiple existing antibiotics, so much so that these bacteria have been classified as ESKAPE and pose a major challenge to clinical anti-infection treatment.²

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The spread of resistance genes is often linked to mobile genetic elements (MGEs), such as plasmids, transposons, and integrons, which are transmitted via horizontal gene transfer (HGT).¹⁰ In *P. aeruginosa*, resistance genes move to recipient bacteria by the HGT of plasmids, integrative and mobilizable elements (IMEs), or integrative and conjugative elements (ICEs).^{11,12} Conjugative plasmids encode products for facilitating conjugation, including relaxase (induction of DNA strand breaks and rejoining), type IV secretion systems (transmembrane channel linking donor and recipient bacteria and DNA transfer), *oriT* (origin of transfer), and coupling proteins (intracellular machinery to link the secretion system).¹³ IMEs typically utilize conjugation mechanisms encoded by other MGEs to achieve transfer and integration.¹⁴ while ICEs mainly use tyrosine/serine recombinases or DDE transposases for chromosomal integration and excision.¹⁵ In previous study, a number of novel MGEs were identified in *P. aeruginosa* in China, such as the novel Inc groups: Inc_{p60512–IMP} and Inc_{pPA7790};¹⁶ Tn6852-related IMEs carrying *bla*_{PER-1} and *bla*_{CARB-2}: Tn6853;¹⁷ Tn6417-related ICEs carrying *bla*_{VIM-1}: Tn6953;¹⁸ and Multiple Tn1403-related transposons carrying different class 1 integrons: Tn6847, Tn6849, Tn7454, Tn7484, etc.^{16,17,19} Multiple types of novel MGEs are frequently identified in China, which pose unknown challenges to the containment of bacterial resistance. It is necessary to pay close attention to the emergence of novel MGEs and their genetic evolution characteristics.

This study aims to describe the genetic structure and evolutionary characteristics of novel MEGs to explore the evolution and role of MGEs in the transmission of antibiotic resistance in *P. aeruginosa*. We sequenced 15 *P. aeruginosa* isolates from China. Detailed genetic dissection and comparison were applied to 19 MGEs, including 15 identified here and an additional four obtained from GenBank. These 19 MGEs were classified into seven types: $Inc_{pSE5381-aadB}$ plasmids; Tn7495-related IMEs; Tn7665-related IMEs; Tn6417-related ICEs; IS-based transposition units; and two unit transposons. In addition, we also identified three novel variants of class D β -lactamases and characterized their antibiotic resistance phenotypes. These findings provide a deeper understanding of the genetic characteristics and diversity of MGEs in *P. aeruginosa*.

Materials and Methods

Bacterial Strains

In the initial stage of this study, more than 1000 drug-resistant *P. aeruginosa* bacteria were isolated from clinical samples in China for large-scale next-generation sequencing (NGS). After preliminary prediction and screening, the isolates carrying multiple ARGs or novel MGEs were screened for third-generation sequencing (TGS). Finally, 15 clinical isolates of *P. aeruginosa* carrying novel MGEs were included in this study. These fifteen *P. aeruginosa* isolates (Table S1) were isolated from various fluid specimens from patients in five different Chinese public hospitals from 2011 to 2018. Bacterial species were identified using average nucleotide identity analysis based on genome sequencing (http://www.ezbiocloud.net/tools/ani).²⁰ Antimicrobial susceptibility testing was performed using VITEK 2 (Table S2) and interpreted according to the 2022 Clinical and Laboratory Standards Institute (CLSI) guidelines.²¹

Sequencing and Sequence Assembly

Bacterial genomic DNA was extracted using the UltraClean Microbial Kit (Qiagen, NW, Germany) and sequenced from a sheared DNA library with the average size of 15 kb (range: 10 to 20 kb) on a PacBio RSII sequencer (Pacific Biosciences, CA, USA). A paired-end library with an average insert size of 350 bp (range: 150 to 600 bp) was sequenced

on a HiSeq sequencer (Illumina, CA, USA). Paired-end short Illumina reads were used to correct long PacBio reads with proovread,²² corrected reads were assembled using *SMARTdenovo* (<u>https://github.com/ruanjue/smartdenovo</u>). As mentioned earlier, further sequence data mining was carried out.^{23,24}

Multi-locus Sequence Typing (MLST)

Sequence types (STs) of *P. aeruginosa* isolates were identified according to the online *P. aeruginosa* MLST scheme (<u>https://pubmlst.org/paeruginosa/</u>). This method mainly performs sequence typing by comparing the diversity of the core sequences of multiple housekeeping genes.

Sequence Annotation and Comparison

Open reading frames (ORF) and pseudogenes were predicted using *RAST 2.0*²⁵ combined with *BLASTP/BLASTN* searches²⁶ against the *RefSeq*²⁷ and *UniProtKB/Swiss-Prot* databases.²⁸ Resistance genes, MGEs, and their characteristics were annotated using online databases including *ResFinder*,²⁹ *Tn Number Registry*,³⁰ *ISfinder*,³¹ *INTEGRALL*,³² and *CARD*,³³ as well as our published database, DANMEL.³⁴ Multiple and pairwise comparisons were performed using *BLASTN/BLASTX*. Gene organization diagrams were created using Inkscape 1.0 (https://inkscape.org/en/).

Cloning Experiments

Coding regions of $bla_{OXA-1202/1203/1204}$ together with 443-bp upstream (promoter) sequence and 160-bp downstream (terminator) sequence from isolates NY7570/SE5381/NY11384, respectively, were subcloned into the vector plasmid pUC57-Kan. In parallel, coding regions for the other bla_{OXA} variants together with the above promoter- and terminator-proximal region were synthesized and subcloned into pUC57-Kan. Resulting recombinant plasmids were transformed into *E. coli* TOP10 by electroporation to generate the corresponding electroporant. 200mg/mL ampicillin (for bla_{OXA} . $_{1202/1203/1204}$) was used for electroporant selection and antimicrobial susceptibility testing was performed using the classic broth microdilution method and interpreted according to the 2022 Clinical and Laboratory Standards Institute (CLSI) guidelines.²¹

Nucleotide Sequence Accession Numbers

The complete chromosome sequences of SE5356, SE5416, NY11084, NY7723, NY5051, NY11352, SE5430, SE5431, NY7570, NY4383, NY3649, NY7583, SE5381, NY11384, and SE5418 and those plasmids of pNY7583-catB3, pSE5381-aadB, pNY11384-aadB, and pSE5418-aacA3 were submitted to GenBank under accession numbers CP120704, CP046404, CP120705, CP120708, CP120707, CP132352, CP054790, CP054791, CP120857, CP120859, CP124612, CP120881, CP054789, CP120883, CP054790, CP120882, CP121188, CP120884, CP123896, respectively.

Results

Identification of P. Aeruginosa Isolates

Identification of Seven STs and T3SS Virulotypes from the 15 Clinical P. Aeruginosa Isolates

Seven STs were identified from the 15 sequenced *P. aeruginosa* isolates, including ST235, ST244, ST292, ST606, ST697, ST740, and ST1284 (<u>Table S1</u>). Among these, eight belonged to ST235 or ST244, considered among the global top 10 *P. aeruginosa* high-risk clones. Four isolates, NY7583, SE5381, NY11384, and SE5418 belonged to ST244; four others, NY5051, NY11352, SE5430, and NY4383 belonged to ST235. Three isolates, SE5356, SE5416, and NY11084 belonged to ST697. Isolates NY7723, SE5431, NY7570, and NY3649 belonged to ST606, ST1284, ST740, and ST292, respectively.

The type 3 secretion system (T3SS) is a virulence system used by *P. aeruginosa* and exhibits widespread presence within the species. T3SS-associated virulence genes (*exoT*, *exoY*, *exoS*, and *exoU*) were identified through BLAST searches and PCR analysis (Table S1). Among the 15 isolates, *exoT* was detected in all (n=15), followed by *exoY* (n=13), *exoS* (n=9) and *exoU* (n=5). The most common virulotype was exoT+/exoY+/exoS+/exoU-.

Identification of 15 MGEs from the 15 P. Aeruginosa Isolates

Whole genome sequencing was performed on 15 *P. aeruginosa* isolates, leading to identification of 15 MGEs that subsequently underwent detailed genetic characteristic analysis: i) pNY7583-catB3, pSE5381-aadB, pNY11384-aadB, and pSE5418-aacA3 from isolates NY7583, SE5381, NY11384, and SE5418, respectively; ii) Tn7495, Tn7496a, and Tn7496b from isolates SE5356, SE5416, and NY11084, respectively; and iii) Tn7666, Tn7667, and T7665RE_{NY11352} from isolates NY7723, NY5051, and NY11352, respectively; iv) Tn7671 from isolates NY7570; v) Tn7669 and Tn7670 from isolates SE5430 and SE5431, respectively; vi) Tn7672, Tn7673 from isolates NY4383, NY3649, respectively (Table 1).

Group	MGE	Accession number	Host Bacterium	Nucleotide Position	Length (bp)	Reference
Tn7495-related IMEs	Tn7495	CP120704	P. aeruginosa SE5356	Chromosome (6,187,966.6237466)	49,501	This study
	Tn7496a	CP046404	P. aeruginosa SE5416	Chromosome (6,367,612.6417112)	49,501	This study
	Tn7496b	CP120705	P. aeruginosa NY11084	Chromosome (6,448,423.6499534)	51,112	This study
Tn7665-related elements	Tn7665	EU595750	P. aeruginosa PACS171b clone fa1389	ginosa PACS171b Not applicable		[35]
	Tn7666	CP120708	P. aeruginosa NY7723	Chromosome (2,756,790.2792449)	35,660	This study
	Tn7667	CP120707	P. aeruginosa NY5051	Chromosome (2,473, 150.253 1255)	58,106	This study
	T7665RE _{NY11352} - I	CP132352	P. aeruginosa NY11352	Chromosome (4,775,980.4852478)	76,499	This study
IS222-based transposition unit	Tn7669	CP054790	P. aeruginosa SE5430	Chromosome (2,299,605.2335185)	35,581	This study
IS6100-based transposition unit	Tn7670	CP054791	P. aeruginosa SE5431	Chromosome (3,375,085.3397228)	22,144	This study
Tn6417-related ICEs	Tn6417	CP013993	P. aeruginosa DHSOI	Chromosome (5,365,108.5473293)	108,186	[36]
	Tn7671	CP120857	P. aeruginosa NY7570	Chromosome (4,960,997.5090637)	129,641	This study
Tn501-related unit transposons	Tn <i>501</i>	Z00027	P. aeruginosa PAT	Plasmid pVSI (Not applicable)	8355	[37]
	Tn7672	CP120859	P. aeruginosa NY4383	Chromosome (344,468.361266)	16,799	This study
Tn <i>1403</i> -related unit transposons	Tn <i>l 403</i>	AF313472	P. aeruginosa RPLII	RPLII Plasmid RPLII 19,630 (Not applicable)		[38]
	Tn7673	CP124612	P. aeruginosa NY3649	Chromosome (3,343,627.3364829)	21,203	This study

Table I Major Features of MGEs Characterized in This Work

(Continued)

Group	MGE	Accession number	Host Bacterium	Nucleotide Position	Length (bp)	Reference
Inc _{pSE5381-aadB}	pNY7583-catB3	CP120882	P. aeruginosa NY7583	-	28,293	This study
	pSE5381-aadB	CP121188	P. aeruginosa SE5381	-	26,620	This study
	pNY11384-aadB	CP120884	P. aeruginosa NY11384	-	26,618	This study
	pSE5418-aacA3	CP123896	P. aeruginosa SE5418	-	25,355	This study

Collection of 19 MGEs for Detailed Sequence Comparison

A detailed genome sequence comparison was applied to a total of 19 MGEs (14 chromosome-borne MGEs and four reference/ prototype MGEs from GenBank: Tn7665,³⁵ Tn6417,³⁹ Tn501,⁴⁰ and $Tn1403^{41}$) that were classified into seven types, which were divided into three groups in this study (Table 1): i) novel families of MGEs (firstly identified in this study): $Inc_{pSE5381-aadB}$ plasmids (including pNY7583-catB3, pSE5381-aadB, pNY11384-aadB, pSE5418-aacA3) and Tn7495-related IMEs (including Tn7495, Tn7496a, and Tn7496b); ii) newly defined MGEs (firstly designated in this study, but with previously determined sequences): Tn7665-related IMEs (including Tn7665, Tn7666, Tn7667, and its derivatives $T7665RE_{NY11352}$); iii) novel transposons with reference prototypes identified in this study: Tn6417-related ICEs (including Tn6417 and Tn7671), IS-based transposition units (including Tn7669 and Tn7670), Tn501-related unit transposons (including Tn7672), Tn1403related unit transposons (including Tn1403 and Tn7673). In addition, 12 novel class 1 integrons were identified in these 19 MGEs. At least 36 antibiotic resistance genes (ARGs), involved in resistance to 11 different categories of antimicrobials and heavy metals, were identified among these 19 MGEs (Figure 1 and <u>Table S3</u>).



Figure I Heatmap of prevalence of resistance genes. Note: The original data are shown in <u>Table S3</u>.

Novel Families of MGEs Identified in This Study

A Novel Group of Inc_{pSE5381-aadB} Plasmids

We characterized four novel plasmids: pNY7583-catB3, pSE5381-aadB, pNY11384-aadB, and pSE5418-aacA3. All had a *rep* (replication initiation protein) gene that could not be assigned to any known Inc groups, so were proposed as a novel Inc group, $Inc_{pSE5381-aadB}$. A detailed sequence comparison of these four plasmids (Figure 2) revealed a highly similar backbone structure (nucleotide identity >90%), including *rep*, *parA/E*, *tivB6/8/10*.

Each of pNY7583-catB3, pSE5381-aadB, pNY11384-aadB, and pSE5418-aacA3 contained two accessory modules. First, each of the four plasmids acquired a novel concise class 1 integron In4875 (see below), In1797 (see below), In4876 (see below), and In1798 (see below), respectively, integrated at the same site downstream of *copG*. Moreover, pNY7583-catB3, pSE5381-aadB, and pNY11384-aadB harbored a complete Tn*5393c* at the same site downstream of *orf210*, while pSE5418-aacA3 had a Δ Tn*5393c* interrupted by IS6100. There was also a major modular difference, which occurred only in pSE5418-aacA3, involving a 9.5 kb inversion likely caused by the insertion of IS6100 into Tn*5393c*.

Three Related Novel IMEs Tn7495, Tn7496a, and Tn7496b

Tn7495 (49.5 kb in length), Tn7496a (49.5 kb in length), and Tn7496b (51.1 kb in length) were integrated at the same site upstream of the *P. aeruginosa* chromosomal gene tRNA^{Ser}. These three IMEs had a highly similar backbone, including core backbone markers *attL*, *attR*, *int1*, and *int2* (Figure 3a). Tn7495 differed from Tn7496 only in the insertion of integrons: In1796 (see below) and In2094 (see below), integrated at the same site within Tn7495 and Tn7496. In comparison to Tn7496a, Tn7496b inserted an ISC*fr1* at the IRt (inverted repeat at the tni end) site of integron In2094.



Figure 2 Linear comparison of the four $Inc_{pSE5381-aadB}$ plasmids.

Notes: Genes are denoted by arrows. Genes, MGEs, and other features are colored based on their functional classification. Shading denotes regions of homology (nucleotide identity \geq 90%).



Figure 3 Comparison of three Tn7495 related IMEs and five Tn7665 related elements. Notes: (a) Comparison of three Tn7495-related IMEs; (b) Comparison of five Tn7665-related elements. Genes are denoted by arrows. Genes, MGEs, and other features are colored based on their functional classification. Shading denotes regions of homology (nucleotide identity \geq 90%). Numbers in brackets indicate nucleotide positions within the chromosome of isolates SE5356, SE5416, NY11084, PACS171b, NY7723, NY5051, and NY11352. The accession number of Tn7665 used as reference is EU595750.³⁵

Newly Defined MGEs Identified in This Study

Tn7665 and Its Derivatives Tn7666, Tn7667, T7665RE_{NY11352}-1, and T7665RE_{NY11352}-2

Tn7665(PACS171b GI),³⁵ a prototype IME originally discovered from *P. aeruginosa* PACS171b, had a 33.0 kb backbone (containing *int1, int2*, and a tellurite resistance operon region). Tn7665, Tn7666, Tn7667, and T7665RE_{NY11352} were integrated at the same site of the *P. aeruginosa* chromosomal gene *endA* (DNA-specific endonuclease I) and producing 2-bp direct repeats (5'-CA) (Figure 2b) Their backbones had at least two main modular differences: i) the *orf1260*–to–*orf303* region was found only in Tn7665 and Tn7666; ii) Tn7667 and T7665RE_{NY11352} had similar backbones; however, T7665RE_{NY11352} was divided into T7665RE_{NY11352}-1 and T7665RE_{NY11352}-2 (found downstream of T7665RE_{NY11352}-1) and the *orf1215*-to-*orf303* region was absent in the backbone structure of T7665RE_{NY11352}.

No accessory modules were detected in Tn7665, whereas Tn7666, Tn7667, and T7665RE_{NY11352} exhibited distinct accessory modules (Figure 3b): i) Tn7666 acquired a single accessory module IS*Pst3*, inserted into the site of *orf1401*; ii) Tn7667 and T7665RE_{NY11352}-1 acquired a Tn*1403* related unit transposon Tn7668 (see below) and Tn7497 (see below), respectively, integrated into the same site of *orf1206*; iii) the *orf1215* gene of T7665RE_{NY11352}-1 and the *orf1401* gene of T7665RE_{NY11352}-2 each harbored an IS26 element. Notably, we found a Tn6582-related ICE at the upstream site of T7665RE_{NY11352}-2, which may have caused the interruption or movement of T7665RE_{NY11352} due to its identification of the IS26 site.

Novel Transposons with Reference Prototypes Identified in This Study Comparison of Two Related ICEs Tn6417 and Tn7671

Tn6417,³⁶ a 108.2 kb ICE first found in *P. aeruginosa* DHS01,³⁹ had an 86.0 kb backbone (containing *attL*/R, *int, cpl*, and *rlx*) and an insertion module, Tn6532. Tn7671 (129.6 kb in length) was identified from *P. aeruginosa* NY7570. Tn6417 and Tn7671 were integrated at the same site downstream of the *P. aeruginosa* chromosomal gene tRNA^{Gly}. There were three main modular differences in their backbones: i) Tn6417 contained its unique backbone region *orf1371, smc*-to-*hepA* region, *zot*-to-*orf249* region, and *orf378*-to-*orf1068* region; ii) Tn7671 harbored a unique backbone region including *orf828, orf306, piL1*-to-*orf189* region, *orf3336*-to-*orf1758* region, *zot*-to-*orf219* region, and *orf378*-to-*orf2514* region from Tn7671 shared <90% nucleotide identity with the corresponding regions of Tn6417 (Figure 4). Like Tn6417, Tn7671 also carried a single accessory module In4877 (see below), a novel concise class 1 integron, which was integrated into the gene *ftsk*, which encodes a cell division protein.

Four Transposons Tn7669, Tn7670, Tn501, and Tn7672

Tn7669 and Tn7670 (Figure 5a and b) belong to the IS-based transposition units and underwent mobilization as a complete unit, bordered by IRL/IRR. Tn7669 is an IS222-based transposition unit, and harbors an IS222 together with captured *fosA* genes (fosfomycin resistance) and a novel concise class 1 integron In1800 (see below). Tn7669 is an IS6100-based transposition unit, and harbors IS6100 together with captured 18.1 kb T1403RE_{SE5431}-1 (see below) and ISCR1.



Figure 4 Comparison of two related ICEs Tn6417 and Tn7671.

Notes: Genes are denoted by arrows. Genes, MGEs and other features are colored based on their functional classification. Shading in light blue denotes regions of homology (nucleotide identity \geq 90%), light orange (nucleotide identity \leq 90%). Numbers in brackets indicate nucleotide positions within the chromosome of isolate NY7570. The accession number of Tn6417³⁹ used as reference is EU696790.



Figure 5 Organization of Tn7669, Tn7670, Tn501 and Tn7672.

Notes: (a) Organization of Tn7669; (b) Organization of Tn7670; (c) Comparison of Tn501 and Tn7672. Genes are denoted by arrows. Genes, MGEs, and other features are colored based on their functional classification. Shading denotes regions of homology (nucleotide identity \geq 90%). Numbers in brackets indicate nucleotide positions within the chromosome of isolates SE5630, SE5431, and NY4383. The accession number of Tn501⁴⁰ used as reference is Z00027.

Tn501 and Tn7672 belong to the Tn3-family unit transposons. The prototype transposon Tn501 was initially discovered in pVS1, a non-conjugative plasmid carried by *P. aeruginosa* PAT.^{37,42} Tn501 has a 12.6 kb backbone with the core backbone structure IRL–*tnpA*–*tnpR*–*res*–*mer*–IRR. Tn7672 (16.7 kb in length) has almost all Tn501 core backbone structure (Figure 5c), and showed two main modular differences: i) Tn7672 contained its unique backbone region *orf825* and *orf1560*; ii) Tn501 carried no accessory modules, but Tn7672 acquired a novel class 1 integron In1825 (see below) and IS6100.

Tn/403 and Its Derivatives Tn7673, Tn7668, Tn7497, TI403RE_{SE5431}-1, and TI403RE_{SE5431}-2

A sequence comparison (Figure 6) was conducted on five Tn1403 derivatives, namely Tn7673, Tn7668, Tn7497, T1403RE_{SE5431}-1, and T1403RE_{SE5431}-2 (Tn7668, Tn7497, T1403RE_{SE5431}-1, and T1403RE_{SE5431}-2 were identified from the above-mentioned elements Tn7667, Tn6861, and Tn6876, respectively), together with the GenBank-derived reference sequence Tn1403. Tn1403³⁸ (19.6 kb in length), a prototypal unit transposons of the Tn3-family, initially identified in plasmid RPL11 from *P. aeruginosa*, harbored a core backbone structure IRL-*tnpA*-*tnpR*-*res*-*sup*-*uspA*-



Figure 6 Comparison of Tn1403 and five related elements.

Notes: Genes are denoted by arrows. Genes, MGEs and other features are colored based on their functional classification. Shading in light blue denotes regions of homology (nucleotide identity \geq 90%). Numbers in brackets indicate nucleotide positions within the chromosome of isolates SE5431, NY3649, NY5051, and NY11352. The accession number of Tn/403⁴¹ used as reference is AF313472.

dksA–yjiK–IRR. Tn7673, Tn7668, and Tn7497 shared the same core backbone structure as Tn1403, while Tn7668 and Tn7497 contained a unique backbone region *crp*–to–*orf693* region bracketed by 9-bp direct repeats (5'-CCCATGACG). T1403RE_{SE5431}-1, and T1403RE_{SE5431}-2 were recognized as two parts of a disrupted MGE.

These Tn1403 derivatives' backbones had at least three major modular differences. First, each acquired one or more integrons into the *res* site, instead of In28 (GCA: bla_{CARB-2} -*cmlA1d*-*aadA1*) in Tn1403: class 1 integrons In1801 (see below), In1829 (see below), and In2252 (see below) were inserted in T1403RE_{SE5431}-1, Tn7673, and Tn7668, respectively; two In51 were integrated into Tn7497. Second, Tn1403, Tn7673, and T1403RE_{SE5431}-2 acquired a Tn5393c at the same site of backbone gene *uspA*; however, integron In1829 of Tn7673 inserted an IS6100, which might lead to disruption of Tn5393c and inversion of a 6.3 kb region (including IS6100, backbone region $\Delta res-3'$ -*sup*- $\Delta uspA-5'$ and disrupted Tn5393c-5' fragment). Third, Tn7668 and Tn7497 carried a Δ Tn6758 (*mer* region from Tn6758) at the same site downstream of *crp*-to-*orf*693 region, and Tn7497 also harbored a unique 20.8 kb MDR region composed of IS*CR1*-*armA* unit, IS*Ec29*-*mph*(E)-IS26 unit, and In27. Moreover, insertion of a 20.8 kb MDR region might be responsible for a 22.2 kb inversion in Tn7497 (including In51, backbone region $\Delta res-3'$ -to-*orf*693, and Δ Tn6758).

Comparison of 12 Novel Integrons

Twelve novel integrons were discovered among the 15 newly sequenced MGEs; all were concise class 1 integrons (Figure 7). Except for In1801 (GCA: *aacA3–aadA2–cmlA1b–catB11*), which is truncated, all remaining 11 integrons exhibited the core integron structure (including IRi/IRt, 5'-CS, GCA, 3'-CS, and *tin*₄₀₂ module). Among them, In1800 (GCA: *aadB–aacA4'–aadB*)



Figure 7 Comparison of the 12 novel integrons.

Notes: Genes are denoted by arrows. Genes, mobile elements and other features are colored based on their functional classification. Shading denotes regions of homology (nucleotide identity \geq 90%). Numbers in brackets indicate nucleotide positions within the corresponding plasmids pSE5381-aadB, pSE5418-aacA3, pNY7583-catB3, and pNY11384-aadB or within the chromosomes of isolates NY3649, NY4383, SE5430, NY5051, SE5431, SE5356, SE5416, and NY7570.

contained a complete *tniABQR* module, enabling self-transfer and facilitating enhanced mobility of mobile components when integrated into elements such as transposons. On the other hand, In2252 (GCA: bla_{VIM-4} – bla_{VIM-4} –aacA27–aacA29f) contained a truncated *tniABQR* module, potentially resulting in impaired its self-transferability.

In addition, except for In1798 (GCA: aacA3-catB3), In1825 (GCA: aacA4'-aadB), and In1800, the other nine integrons had at least one insertion module in the 5'-CS or 3'-CS or tni_{Tn402} : i) In1797 (GCA: $aadB-bla_{OXA1203}$), In4875 (GCA: $aacA4'-bla_{OXA-101}$), In4876 (GCA: $aadB-bla_{OXA-1204}-aphA15-catB3$) and In1829 (GCA: $bla_{OXA-2}-aacA4-guc207-bla_{OXA-2}$) exhibited the insertion of one IS6100 at the site downstream of the 3'-CS region; ii) In2252 was inserted by a separate module ISPa7 in the 5'-CS region; iii) In1801 (GCA: aacA3-aadA2-cmlA1b-catB11) had In0 and ISCR3-tetA(G)-cmlA9 unit; (iv) In1796 (GCA: cmlA1b-aadA1dy) and In2094 (GCA: cmlA1b-aadA1a) harbored In0, ISCR3-tetA(G)-cmLA9 unit, and Δ Tn6758; (v) In4877 (GCA: $bla_{OXA-1202}-aacA4-crf240-bla_{CARB-2}$) acquired an aadA2-related region at the site of $\Delta intI1$ and an IS6100 at the site downstream of orf6, which was interrupted by a complete Tn6758. In particular, $bla_{OXA-101}$ and its three novel variants $bla_{OXA-1202/1203/1204}$ were identified within the GCA of integrons In4875, In4877, In1797, and In4876, respectively.

Summary of Newly Identified or Designated MGEs

We identified 23 new MGEs, including four IMEs Tn7495, Tn7496, Tn7666, and Tn7667; one ICE Tn7671; four unit transposons Tn7668, Tn7497, Tn7672, and Tn7673; two IS-based transposition units Tn7669 and Tn7670; and 12 integrons In1796, In2094, In2252, In4877, In1825, In1800, In1801, In1829, In4875, In1797, In4876, and In1798. One novel Inc group Inc_{pSE5381-aadB} was designated. Additionally, the element PACS171b GI previously designated with determined sequences was renamed as standard Tn designations Tn7665.

Three Novel Variants of Antibiotic Resistance Genes and Their Characteristics

Three novel $bla_{OXA-101}$ variants encoding OXA- β -lactamase, namely $bla_{OXA-1202}$, $bla_{OXA-1203}$, and $bla_{OXA-1204}$ (Figure S1), were identified from three distinct clinical isolates: *P. aeruginosa* NY7570, SE5381, and NY11384, respectively. These three novel variants, all of which were categorized within the OXA-10-like OXA enzyme, featured genes that were 801 bp in length and encoded a 266 amino acid-long protein. The deduced OXA-1202 differs from OXA-101 by a single amino acid substitution Val188Ala; OXA-1203 differed from OXA-101 by one amino acid substitution Ala66Val; OXA-1204 differed from the OXA-101 by one amino acid substitution Trp154Gly.

OXA-1202/1203/1203 gene fragments, their point mutants OXA-101, and the prototype OXA variant OXA-10 (as a positive control) were cloned into pUC57K (a kanamycin-resistant plasmid) and transformed into *E. coli* TOP10 to obtain *E. coli* electroporants TOP10/pUC57K-OXA-1202/1203/1204/101/10. Classic broth microdilution was applied to test antimicrobial susceptibility of these electroporants and their negative control strains (TOP10, TOP10/pUC57K, and ATCC 25,922). Strains encoding OXA-1202/1203/1204 enzymes were highly resistant to amoxicillin and ampicillin relative to the negative controls, but were still susceptible to piperacillin (Table 2). Furthermore, TOP10/pUC57K-OXA-1204 was also resistant to ceftazidime and cefoxitin, while TOP10/pUC57K-OXA-1202/1203 was susceptible to cephalosporins and exhibited narrow-spectrum antibiotic resistance similar to OXA-101.

Bacterial Isolate	Minimum Inhibitory Concentration ($\mu g/mL$)/Antimicrobial Susceptibility						
	Amoxicillin	Ampicillin	Piperacillin	Cefazolin	Cefuroxime	Ceftazidime	Cefoxitin
TOP10/pUC57K-OXA-10	>1024	>1024	256	8	16	0.25	16
TOP10/pUC57K-OXA-101	>1024	512	128	4	8	0.25	8
TOP10/pUC57K-OXA-1202	>1024	512	32	4	16	0.5	8
TOP10/pUC57K-OXA-1203	>1024	512	32	4	16	0.5	8

Table 2 MICs of Selected Antimicrobial Agents for the Transformants Carrying $\mathsf{bla}_\mathsf{OXA}$

(Continued)

Bacterial Isolate	Minimum Inhibitory Concentration (µg/mL)/Antimicrobial Susceptibility							
	Amoxicillin	Ampicillin	Piperacillin	Cefazolin	Cefuroxime	Ceftazidime	Cefoxitin	
TOP10/pUC57K-OXA-1204	>1024	128	8	4	16	16	32	
TOP10/pUC57K	16	8	8	4	8	0.25	8	
TOP10	16	8	8	4	8	0.25	8	
ATCC 25922	8	8	4	4	8	0.25	4	

Table 2 (Continued).

Discussion

Fifteen multiple drug resistance (MDR) isolates were isolated from clinical samples and completely sequenced in this study. Among these isolates, ST235 and ST244 are the more common high-risk clones. All ST335 isolates harbored *exoU* +/*exoS*- virulotypes, while all ST244 isolates harbored *exoU*-/*exoS*+ virulotypes. This finding indicates a potential correlation between the high-risk clones and T3SS virulotypes, consistent with previous work.^{43,44} In addition, T3SS virulence genes were detected in all 15 MDR isolates collected. Previous studies have shown a correlation between the presence of virulence genes and bacterial resistance, and antibiotic resistance genes has been easily detected in T3SS-positive isolates.^{45,46} which might jointly contribute to the high prevalence of *P. aeruginosa*.

A total of 15 novel MGEs were identified through comprehensive genome alignment. In particular, the $Inc_{pSE5381-aadB}$ plasmids and Tn7495-related IMEs were identified and sequenced for the first time. By blasting the rep sequence and the backbone sequence among GenBank (identity \geq 95%, last accessed 20 Feb, 2023), we found that $Inc_{pSE5381-aadB}$ plasmid was only harbored in *P. aeruginosa*. To verify its transfer ability, we performed multiple conjugation experiments, but did not succeed in transferring $Inc_{pSE5381-aadB}$ plasmid from their wild isolates to PAO1 or ATCC 27,853, possibly due to incomplete functionality of the plasmid-encoded conjugative transfer protein. Tn7495 family IMEs presented a conserved backbone with the same chromosomal insertion site. Similar to conventional IMEs, Tn7495 family IMEs do not undergo intercellular transfer autonomously. *BLAST* searches of the integrase sequence of Tn7495-related IMEs among GenBank (identity \geq 95%, with relatively complete element backbone, last accessed 20 Feb, 2023) indicated that as of yet these IMEs have only been identified in *P. aeruginosa*. Although the newly discovered elements in this study had only been identified in *P. aeruginosa*, they might also be widely spread to other species in the process of genetic evolution, and this unknown situation deserved our close attention.

As prototypes or references, Tn7665, Tn6417, Tn501, and Tn1403 were all initially found in *P. aeruginosa*. Tn7665related IMEs had been frequently reported in *P. aeruginosa* and there are documented instances of insertions of this group of elements within the chromosomal gene *endA*,⁴⁷ consistent with the findings of this study. Tn6417-related ICEs had been identified not only in *P. aeruginosa* but also occasionally among other species, such as *Klebsiella pneumoniae*, *Morganella morganii*, *Bordetella*, and *Achromobacter xylosoxidans*. The presence of Tn501-related elements and Tn1403-related elements in *P. aeruginosa* has been reported multiple times.^{17,48,49} Overall, the types of MGEs identified in this study exhibit a high prevalence among clinical *P. aeruginosa* isolates, revealing that they may play a critical role in facilitating the spread of resistance in *P. aeruginosa*.

In our study, a total of 36 ARGs were detected from the 19 MGEs analyzed, involving 11 classes of antibiotics. Among them, aminoglycoside resistance genes (n=13) and β -lactam resistance genes (n=7) accounted for the largest proportion. Thirty of these ARGs were found in the gene cassette of class 1 integron; the remaining six belonged to other sub-regions: genes *strA* and *strB* in the Tn5393c; gene *fosA* in the Tn7669 backbone; genes *mph*(E) and *msr*(E) in the IS*Ec29–mph*(E)–IS26 unit; gene *ter* in the Tn7495-related IMEs backbone (Table S3). In addition, we identified three novel β -lactamase genes (*bla*_{OXA-1202/1203/1204}) located within integrons, showing that integron activity not only aided the spread of ARGs but may also expedite the evolution of antibiotic resistance, as found previously.⁵⁰ During the transfer of these MGEs, some mutations or the occurrence of the homologous recombinant events accelerated the evolution of the

MGEs and promoted the production of novel ARGs. These ARGs-carrying MGEs formed intricate mosaic structures and exhibited intercellular or intracellular mobility, facilitating the widespread dissemination of bacterial resistance.

OXA enzymes are a large family of enzymes with more over a thousand variants that are widely distributed. Three newly identified bla_{OXA} variants (bla_{OXA-1202/1203/1204}) in P. aeruginosa belonged to the OXA-10-like subfamily like their point mutant $bla_{OXA-101}$. The presence of OXA-10-like β -lactamases has been repeatedly observed in P. aeruginosa,^{7,51} and has also been occasionally reported in Acinetobacter baumannii⁵² and Enterobacteriaceae.⁵³ According to antibiotic susceptibility testing, OXA-1202, OXA-1203 presented a similar resistance phenotype as their point mutant OXA-101. Notably, OXA-1204 showed reduced hydrolytic activity against some antibiotics tested (eg ampicillin and piperacillin) compared with OXA-101 and exhibited exclusive resistance to ceftazidime and cefoxitin. The substitution of W154G in OXA-1204 might play a role in increasing resistance against certain cephalosporins in P. aeruginosa. Previous studies have shown that lysine carbamylation had important mechanistic functions in the catalytic activity of OXA-10 from P. aeruginosa.^{54,55} The carbamyl group of Lys-70 in OXA-10 can be stabilized by interacting with Ser-67, Trp-154 and a crystal water molecule.⁵⁴ Therefore, the mutation of amino acid 154 in OXA-1204 could potentially influence the carbamylation of Lys-70 and result in a reduction of its hydrolytic activity towards ampicillin and other antibiotics (Figure S2). The diversity of the OXA family continues to expand. While certain inhibitors might impede the activity of these genes' products,⁵⁶ the presence of extended-spectrum class D β-lactamases and carbapenem-hydrolysing class D β -lactamases continues to pose a formidable challenge to clinical anti-infective therapy, necessitating utmost attention.

Conclusion

A detailed sequence comparison of a total of 19 novel MGEs from seven types of *P. aeruginosa* revealed that they carried different combinations of resistance genes or entirely novel resistance gene variants. This study reported the Inc_{pSE5381-aadB} plasmids and Tn7495-related IMEs for the first time, and also identified the novel resistance genes *bla*_{OXA-1202/1203/1204} carried by MGEs. These MGEs underwent continuous modification during transmission, leading to the evolution of intricate mosaic structures through transposition or homologous recombination. In turn, this facilitated the spread of antibiotic resistance and enhanced the adaptability of *P. aeruginosa* under the selective pressure of various antibiotics. The present study has elucidated the genetic evolutionary characteristics of novel ARGs-carrying MGEs, thereby enhancing the diversity of transmission modes of antibiotic resistance in *P. aeruginosa* and providing a deeper understanding of the drug resistance mechanisms of clinical *P. aeruginosa*. In addition, some of the elements identified in this study were only found in *P. aeruginosa*, but these elements are constantly evolving during transmission and may be more widely spread to other species, which deserves our continuous attention in future studies.

Ethics Approval and Informed Consent

This study uses the clinical bacterial isolates obtained from the Chinese public hospitals as listed in <u>Table S1</u>. The local legislation did not require the study to be reviewed or approved by an ethics committee, because the bacterial isolates involved in this study was part of the routine laboratory procedures. The research involving biohazards and all related procedures were approved by the Biosafety Committee of the Beijing Institute of Microbiology and Epidemiology.

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Disclosure

The authors report no conflicts of interest in this work.

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